

Hexaform Hexaloids Uratrine Urodeine



Information on biological activities of small molecules

PubMed Entrez Structure GenBank PubChem Help Search PubChem Compound Compound Summary: O CID: 4101 🛈 🖼 BioActivity: Summary 2 Inactive: 10 Links Inconclusive: 1 Link 🗞 NLM Toxicalagy: 🗵 Link1, Link2 Substances: 🛭 All: 144 Links Same: 39 Links Mixture: 105 Links O Related Compounds: ② Same, Connectivity: 3 Links Similar Compounds: 18 Links (2) Đ, Structure Search 2 MeSH Synonyms **Properties Descriptors** Category Exports O Medical Subject Annotations: (Total:1) 🛛 Methenamine
An anti-infective agent most commonly used in the treatment of urinary tract infections. Its anti-infective action derives from the slow release of formaldehyde by hydrolysis at acidic pH. (From Martindale, The Extra Pharmacopoeia, 30th ed, p173) Show MeSH Tree Structure Pharmacological Action: Anti-Infective Agents, Urinary PubMed via MeSH Choose by Subheadings: administration and dosage adverse effects analogs and derivatives analysis boold chemical synthesis chemistry diagnostic use economics immunology metabolism pharmacokinetics pharmacology poisoning standards therapeutic use therapy toxicity PubMed MeSH Keyword Summary Depositor-Supplied Synonyms: (Total: 123) Display: First 10 | All | Sort: Weight methenamine 🍄 Hexamine 🍄 Urotropin 🍄 Urotropine Uritone Hexamethylenetetramine Antihydral Methenamin Resotropin Aminoform P Ammoform Cystamin Cystogen Duirexol Formamine

Heterin
Hexasan
Formin
Uramin
Aminoformaldehyde
Hexamethylenamine
Preparation AF
Mandelamine
Hexamethylenamine
Hexamethylenamine
Hexamethylenamine
Hexamethylenamine
Methamin
Metramine
Hexa-Flo-Pulver
Ekagom H
Ammonioformaldehyde
Uro-phosphate
Aceto HMT
Herax UTS
Hexamethylentetramin
Nocceler H
Sanceler H
Formin (heterocycle)
Prestwick_79
Hexamethylenetetramine
Hexamine (JAN)
Hexamine (TN)
Hexamine (TN)
Hexamethylenetetramine
Hexamine (USP)
Vulkacit H 30
1,3.5.7-Tetraazadamantane
Hexamethylenetetraminum
HMTA
S 4 (heterocycle)
Hexamethylenetetraminum
HMTA
S 4 (heterocycle)
Hexamethylenetetraminum
HMTA
S 4 (heterocycle)
Hexamethylenetetraminum
HMTA
S 5 (heterocycle)
Hexamethylenetetraminum
HMTA
S 6 (heterocycle)
Hexamethylenetetraminum
HMTA
S 7 (heterocycle)
Hexamethylenetetraminum
HMTA
S 8 (heterocycle)
Hexamethylenetetraminum
HMTA
S 9 (heterocycle)
Hexamethylenetetraminum
HMTA
S 9 (heterocycle)
Hexamethylenetetraminum
HMTA
Horizaminum (INN-Spanish)
Methenamina [INN-Spanish]
Methenamina [INN-Latin]
Esametilentetramina [Italian]
H11300_ALDRICH

Hexamethylenetetramine, tech.
Hexamethylentetramin [German]
KBioGR_001563
KBioSS_001471
DivK1c_000322
SPECTRUM1500394
SPBio_000753
Hexamethylenetetramine solutions
15614_RIEDEL
33233_RIEDEL
CCRIS_2297
CHEB1:6824
H6404_SIAL
HSDB 563
Hexa (vulcanization accelerator)
NSC26346
KBio1_000322
KBio2_001471
KBio2_004039
KBio2_004039
KBio3_00000000
NINDS_000322
NSC403347
AlDS018490
Formin (the heterocyclic compound)
398160_SIAL
AIDS-018490
EINECS_202-905-8
EINECS_246-528-7
NSC_26346
UN1328
EPA Pesticide Chemical Code 045501
NSC 403347
NSC 103-97-0
Al3-09611
Al3,5,7-Tetraazatricyclo[3.3.1.1~3,7~]decane
D00393
Hexamethylenetetramine-palladium chloride adduct
1.3,5,7-Tetraazatricyclo[3.3.1.1,7]decane
1,3,5,7-Tetraazatricyclo[3.3.1.1,7]decane

7

15978-33-3 24911-88-4 56549-34-9 '74734-16-0 91773-48-7

Properties Computed from Structure: 2

Molecular Weight: 140,18628 g/mol Molecular Formula: C₆H₁₇N₄ Hydrogen Bond Donor Count: 0 Hydrogen Bond Acceptor Count: 4 Rotatable Bond Count: 0 Topological Polar Surface Area: 13

\bigcirc Descriptors Computed from Structure: @

Canonical SMILES: C1N2CN3CN1CN(C2)C3
Inch1: Inch1=1/C6H12N4/c1-7-2-9-4-8(1)5-10(3-7)6-9/h1-6H2 ②

Substance Category: 2

Biological Properties: 22 Links
Journal Publishers: 1 Link
Metabolic Pathways: 1 Link
Molecular Libraries Screening Center Network: 2 Links
Physical Properties: 4 Links
Substance Vendors: 6 Links
Theoretical Properties: 1 Link
Toxicology: 3 Links

ASN1 Display Save	XML	Display Save	SDF	Display Save
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PubMed Entrez Structure GenBank PubChem Help Search PubChem Compound Compound Summary: O cid: 407405 🛭 🖽 (BioActivity: Summary (2) Active: 7 Links Inactive: 66 Links NLM Toxicology: Link 🗵 (Substances: 🗓 All: 9 Links Same: 6 Links Mixture: 3 Links Similar Compounds: 5 Links ② (d) Structure Search Đ, Synonymis **Properties** Descriptors Category Depositor-Supplied Synonyms: (Total: 17) 2 Display: First 10 | All | Sort: Weight Allyliodourotropine Allyl iodide hexamine Methenamine allyl iodide Methenamine allyloiodide CBDivE_014100 Iodoallylhexamethylenetetramine Hexamethylenetetramine alliodide NSC5062 NSC7309 Allyl iodide hexamethylenetetramine Hexamethylenetetramine allyl iodide EINECS 253-259-9 NC160_004218 Hexamethylenetetramine, compd. with 3-iodopropene (1:1) Propene, 3-iodo-, compd. with hexamethylenetetramine (1:1) 36895-62-2 3.5, 7-Triaza-1-azoniatricyclo[3.3.1.13,7]decane, 1- (2-propenyl)-, iodide On Properties Computed from Structure: 2 Molecular Weight: 181.25808 g/mol Molecular Formula: C₉H₁₇N₄⁺ Hydrogen Bond Donor Count: 0 Hydrogen Bond Acceptor Count: 3 Rotatable Bond Count: 2 Topological Polar Surface Area: 9.7 ⊕ more... Or Descriptors Computed from Structure: 2 Substance Category: 2 Biological Properties: 4 Links Theoretical Properties: 1 Link Toxicology: 1 Link

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Pub©hem

Information on biological activities of small molecules

PubMed Entrez

HOME SEARCH SITE MAP

Search PubChem Compound GenBank Structure

Compound Summary:

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Parent CID: 5846454 (7) Unique Components: 1 Links CID: 6435993 🗗 🖫

NLM Toxicology: Unk

Substances: 3 Links (2)

Related Compounds: (2)
Same, Connectivity: 2 Links
Similar Compounds: 11 Links (2)

Structure Search D

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Descriptors Properties Synonyms

Category

O Depositor-Supplied Synonyms: (Total: 7)

Sort: Weig

Dowicil 200
Chlorodity methenanine chloride
Chlorodity inectenanine chlorodity chloride, cts isomer
1-(3-Chlorodity)-3,5,7-triaza-1-azmioadamantane chloride, cts form
3,5,7-fnaza-1-azmiatricyclo(3,3,1,13,7)decane, 1-((22)-3-chloro-2-propenyl)-, chloride, 3,5,7-fnaza-1-azmiatricyclo(3,3,1,13,7)decane, 1-(3-chloro-2-propenyl)-, chloride, (2)-51229-78-8

On Properties Computed from Structure:

Molecular Weight: 251.15614 g/mai Molecular Formula: C₂H₁₅C₁P₄ Hydrogen Band Donaro Count: 0 Hydrogen Band Acceptor Count: 1 Hydrogen Bond Acceptor Count: 2 Rotabble Bond Count: 2 Topological Polar Surface Area: 9 /

Descriptors Computed from Structure:

Canonical SMILEs: CINZCHZCHZ(FI+)[C2](C3)(C=CC),[C1-]
Isomerica SMILEs: CINZCHZ(CHZ)(FI+)[C2](C3)(C4)
Inchi: InChi=1/C3H3CHZ(HZ)(C3)(C3)(C4)
Inchi: InChi=1/C3H3CHZ(HZ)(C3)(C3)(C4)
Inchi: InChi=1/C3H3CHZ(HZ)(D4)(D5-1:3-14)-13(1-4)(D3)(1-4

(Substance Category: (2)

Biological Properties: 1 Link Journal Publishers: 1 Link Toxicology: 1 Link

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PubMed Entrez Structure GenBank PubChem Help " Search PubChem Compound Compound Summary: CI O c10: 22730 ☑ ⊕ Parent CID: 22731 ② Unique Components: 2 Links 🔷 NLM Toxicology: Link 📵 Substances: 3 Links 2 Similar Compounds: 3 Links 🖸 Structure Search Ð, Synonyms **Properties** Descriptors Exports Category Openositor-Supplied Synonyms: (Total: 8) Sort: Weight NSC49660 NSC 30209 NSC 49660 1-(2-Thenyl)hexamethylenetetrammonium chloride HEXAMETHYLENETETRAMINIUM, 1-(2-THENYL)-, CHLORIDE 3,5,7-Triaza-1-azoniaadamantane, 1-(2-thienyl)-, chloride Hexamethylenetetramine, compd. with thiophene, 2-(chloromethyl)-6296-08-8 $\begin{array}{lll} \textbf{Molecular Weight: } 272,79748 \ g/mol\\ \textbf{Molecular Formula: } C_{11}\textbf{H}_{17}\text{CIN}_{4}\textbf{S}\\ \textbf{Hydrogen Bond Donor Count: } 0\\ \textbf{Hydrogen Bond Acceptor Count: } 4\\ \textbf{Rotatable Bond Count: } 2\\ \end{array}$ Topological Polar Surface Area: 9.7

more... O Descriptors Computed from Structure: 2 Canonical SMILES: C1N2CN3CN1C[N+](C2)(C3)CC4=CC=CS4.[CI-] InCh1: 1nCh1=1/C11H17N45.CIH/c1-2-11(16-3-1)4-15-8-12-5-13(9-15)7-14(6-12)10-15;/h1-3H,4-10H2;1H/q+1;/p-1/fC11H17N45.CI/h;1h/qm;-1 ②' Substance Category: 🗓 Biological Properties: 2 Links Toxicology: 1 Link





PubMed Entrez Structure GenBank PubChem Help Search PubChem Compound (00 Compound Summary: CID: 64172 1 1 Parent CID: 64173 2 Unique Components: 2 Links NLM Toxicology: Link 🔞 Substances: 2 Links 🗓 Br Similar Compounds: 2 Links 🖸 💢 , Structure Search 🛭 Đ, MeSH Synonyms **Properties** Descriptors Category Exports Medical Subject Annotations: (Total:1) AT 584 PubMed via MeSH PubMed MeSH Keyword Summary Depositor-Supplied Synonyms: (Total: 9) 2 Sort: Weight AT 584 € AT-584 € A1-534 %
-(1-(Bis(2-chloroethyl)amino)phenacyl)-3.5,7-triaza-1-azoniaadamantane bromide
3.5,7-Triaza-1-azoniaadamantane, 1-(p-(bis(2-chloroethyl)amino)phenacyl)-, bromide
Hexamethylenetetramine salt of p-(bis(2-chloroethyl)amino)-alpha-bromoacetophenone
1-(2-(4-(Bis(2-chloroethyl)amino)phenyl)-2-oxoethyl)-3,5,7-triaza-1-azoniatricyclo(3.3.1.1(sup 3,7))decane bromide
16810-17-6 1650-17-6 3.5,7-Triaza-1-azoniatricyclo(3.3.1.1(sup 3,7))decane, 1-(2-(4-(bis(2-chloroethyl)amino)phenyl)-2-oxoethyl)-, bromide 3.5,7-Triaza-1-azoniatricyclo(3.3.1.13,7)decane, 1-(2-(4-(bis(2-chloroethyl)amino)phenyl)-2-oxoethyl)-, bromide Properties Computed from Structure: ② Molecular Weight: 479.24194 g/mol Molecular Formula: C₁₈H₂₆BrCl₂N_SO Hydrogen Bond Donor Count: 0 Hydrogen Bond Acceptor Count: 6 Rotatable Bond Count: 8 Tautomer Count: 2 Topological Polar Surface Area: 30 more... O Descriptors Computed from Structure: 🛽 Canonical SMILES: C1N2CN3CN1C[N+](C2)(C3)CC(=0)C4=CC=C(C=C4)N(CCCl)CCCl,[Br-] InChI: InChI=1/C18H26Cl2N5O.BrH/c19-5-7-24(8-6-20)17-3-1-16(2-4-17)18(26)9-25-13-21-10-22(14-25)12-23(11-21)15-25:/h1-4H,5-15H2:1H/q+1:/ ρ -1/(C18H26Cl2N5O.Br/h;1h/qm;-1 \bigcirc

Substance Category: 2 Biological Properties: 1 Link Toxicalogy: 1 Link

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	Save		Save		Save

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Substance Category: 2

Biological Properties: 4 Links Theoretical Properties: 1 Link

information on biological activities of small molecules Pub(C PubMed Entrez Structure GenBank PubChem Search PubChem Compound Compound Summary: (C) CID: 64160 ☑ ⊞ BioActivity: Summary [2] Inactive: 2 Links 📳 Substances: 🛭 All: 7 Links Same: 5 Links Mixture: 2 Links Similar Compounds: 5 Links (2) Structure Search 2 Đ, Synonyms Properties Descriptors Category Exports Openositor-Supplied Synonyms: (Total: 16) 🗹 Display: First 10 | All | Sort: Weight Formoiodine Hexamethylenetetramine ethyl-Hexamethylenetetramine ethyl iodide NSC7308
NSC7308 (IODINE SALT)
AIDS155858
AIDS-155858
SSC203331
5406-76-8 (IODDINE SALT)
3,5, 7-Triaza-1-azoniaadamantane, 1-ethyl3,5, 7-Triaza-1-azoniaadamantane, 1-ethyl-, iodide
1-Ethyl-Ilambda-5~,3,5,7-tetraazatricyclo[3,3.1.1~3,7~]decane
3,5, 7-Triaza-1-azoniatricyclo[3,3.1.13,7]decane, 1-ethyl3,5, 7-Triaza-1-azoniatricyclo[3,3.1.13,7]decane,) 1-ethyl-, iodide
3,5, 7-Triaza-1-azoniatricyclo[3,3.1.13,7]decane, 1-ethyl-, iodide
5406-76-8 Molecular Weight: 169.24738 g/mol Molecular Formula: $C_BH_{17}N_4^{-4}$ Hydrogen Bond Donor Count: 0 Hydrogen Bond Acceptor Count: 3 Rotatable Bond Count: 1 Topological Polar Surface Area: 9.7 \bigcirc Descriptors Computed from Structure: ②Canonical SMILES: CC[N+]12CN3CN(C1)CN(C3)C2 InChI: InChI=1/C8H17N4/c1-2-12-6-9-3-10(7-12)5-11(4-9)8-12/h2-8H2,1H3/q+1 2

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ASN1	Display	XML	Display	SDF	Display
ASNI	Save	AML	Save	SUF	Save



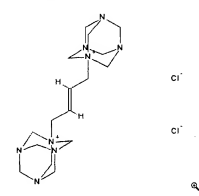
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Structure GenBank Search PubChem Compound

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Help

Compound Summary:



CID: 6435994 ② 9

Parent CID: 6435995 ②
Unique Components: 2 Unks

Substance: 1 Link ②
Similar Compounds: 2 Links ②
. C. Structure Search ②

Synonyms

Properties

Descriptors

Category

Exports

O Depositor-Supplied Synonyms: (Total: 10)

Sort: Weight :-

Cosan 265 Caswell No. 259AA EINECS 257-149-1

EPA Pesticide Chemicol Code 106801
1.1°-(2-Butylene)bis(3,5,7-triaza-1-azonlaadamantane)
1,1°-(2-Butenylene)bis(3,5,7-triaza-1-azonlaadamantane chloride
1,1°-(But-2-en-1,4-diyl)bis(3,5,7-triaza-1-azonlaatricyclo(3.3.1.13,7)decane) dichloride
3,5.7-triaza-1-azonlatricyclo(3.3.1.13,7)decane, 1,1°-(2-butene-1,4-diyl)bis-, dichloride
51350-34-6
59736-58-2

Properties Computed from Structure: 2

Molecular Weight: 405.369 g/mol Molecular Formula: $C_{16}H_{30}Cl_2N_8$ Hydrogen Bond Donor Count: 0 Hydrogen Bond Acceptor Count: 8 Rotatable Bond Count: 4 Topological Polar Surface Area: 19.4 \oplus more...

O Descriptors Computed from Structure: 2

Substance Category: 2

Toxicology: 1 Link

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ASN1	Save	XML	Save	IL	SDF	Save



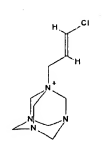
PubMed Entrez Structure

GenBank Search PubChem Compound PubChem

Help 1 (00)

Compound Summary:

CI



O cid: 6433396 ② ₪ Parent CID: 5475987 (2) Unique Components: 2 Links

NLM Toxicology: Link ②

Substances: 3 Links 2 . Poi Related Compounds: 🛭 🕏

Same, Connectivity: 2 Links

Similar Compounds: 11 Links 🛭

(C) Structure Search 2

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MeSH Synonyms

Properties

Descriptors

Category

Exports

Medical Subject Annotations: (Total:1)

quaternium-15

PubMed via MeSH

PubMed MeSH Keyword Summary 2

O Depositor-Supplied Synonyms: (Total: 24)

Display: First 10 | All | Sort: Weight

Dowicide Q Quaternium 15 Quaternium-15 ♥ Dowicil 75 Cinartc 200 Dowicil 100 Caswell No. 181 Dowco 184 CCRIS 1398 HSDB 6820 Methenamine 3-chloroallylochloride EINECS 223-805-0 N-(3-Chloroallyl)hexaminium chloride Pesticide Chemical Code 017901 NSC 172971 XD-1840 XD-1840
Hexamethylenetetramine chloroallyl chloride
N-(3-CHLORALLYL)HEXAMINIUM CHLORIDE
1-(3-Chloroallyl)-3,5,7-triaza-1-azoniaadamantane chloride
3,5,7-Triaza-1-azoniaadamantane, 1-(3-chloroallyl)-, chloride
3,5,7-Triaza-1-azoniatricyclo(3.3.1.13,7)decane, 1-(3-chloro-2-propenyl)-, chloride
103638-29-5 4080-31-3 60789-82-4

 $\mathcal{Q}_{\widehat{\mathbb{Q}}}$ Properties Computed from Structure: 2

Molecular Weight: 251.15614 g/mol Molecular Formula: $C_9H_{16}Cl_2N_4$ Hydrogen Band Donor Count: 0 Hydrogen Bond Acceptor Count: 4
Rotatable Bond Count: 2
Topological Polar Surface Area: 9.7

more...

Substance Category: 2

Biological Properties: 1 Link Journal Publishers: 1 Link Toxicology: 1 Link

ASN1	Display	XML	Display	SDF	Display
ASNI	Save	AML	Save	305	Save

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